

Arney 10-18-4
Serial No. 10/798,064

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CENTRAL FAX CENTER

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Claims Listing

1. (Currently Amended) An implantable stent comprising:
a tubular member having an interior surface and an exterior surface, characterized in that
at least one of said surfaces is being hydrophobic, and
5 a region of said at least one surface includes an array of microstructures or nanostructures
that coverings first portions of said surface, said array causing the region to have a dynamically
controllable hydrophobicity.

2. (Original) The stent of claim 1, further including a control device affixed to said
10 tubular member for varying said hydrophobicity.

3. (Original) The stent of claim 2, wherein said control device comprises an
electronic device or an optical device.

15 4. (Original) The stent of claim 3, wherein said control device is remotely actuatable
from an external source.

5. (Original) The stent of claim 1, wherein said array leaves second portions of said
surface exposed, and further including a chemically active substance adhered to at least one of
20 said exposed second portions.

6. (Original) The stent of claim 5, wherein said substance comprises a
pharmacological agent or a drug.

25 7. (Original) The stent of claim 6, further including a control device affixed to said
tubular member, said device being capable of releasing said agent or drug from said at least one
second portion.

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8. (Original) The stent of claim 7, further including
an electrically conductive substrate that is configured to be electrically isolated from
body fluid in contact with said array of microstructures or nanostructures, and
wherein said control device is capable of applying a voltage between said array and said
5 substrate to vary the penetration of the interstices of said array by said fluid, thereby causing
release of said agent or drug into said fluid.

9. (Original) The stent of claim 1, wherein said array leaves second portions of said
surface exposed, and further including
10 means for electrically isolating said array into separate spatial zones,
at least two of said zones containing chemically active substances adhered to the exposed
second portions thereof, and
wherein said control device is capable of causing the release of said substances of the
separate zones at different times.

15 10. (Original) The stent of claim 9, wherein said substances are the same chemically
active substances of the same or a different dose.

20 11. (Original) The stent of claim 9, wherein said substances are different chemically
active substances.

12. (Original) The stent of claim 1, further including means for altering the shape of
said stent *in vivo*.

25 13. (Original) The stent of claim 12, wherein said altering means is capable of
changing the diameter of said tubular member.

14. (Original) The stent of claim 1, wherein said tubular member has an elongated
slot that is coextensive with its length, thereby forming a pair of elongated edges that are

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movable relative to one another, and the stent further comprising a plurality of electrically controllable structures thereon, the structures capable of moving said edges and releasably latching said edges.

5 15. (Original) The stent of claim 1, wherein said tubular member comprises a semiconductor substrate and said array of microstructures or nanostructures is disposed on said substrate.

10 16. (Original) The stent of claim 15, wherein said tubular member further comprises a layer disposed on said substrate, said substrate and said layer having different thermal expansion coefficients.

 17. (Original) The stent of claim 16, wherein said microstructures or nanostructures have at least one dimension that is in the range of 4 μm to 20 nm.

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 18. (Currently Amended) An implantable stent comprising
a tubular member including a conducting substrate, said member having an interior
surface and an exterior surface, ~~characterized in that~~
at least one of said surfaces is being hydrophobic to a body fluid, and
20 a region of said at least one surface including an array of microstructures or
nanostructures that covers first portions of said surface, said array rendering the region to have a
dynamically controllable hydrophobicity,
a medicinal substance adhered to an exposed second portion of said surface, and
a control device affixed to said tubular member for applying a voltage between said fluid
25 and said substrate to vary said hydrophobicity and release said substance into said body fluid,
said device being actuatable from an *ex vivo* source.

 19. (Original) The stent of claim 18, wherein
said exposed second portion is electrically isolated into first and second spatial zones,

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each zone containing a medicinal substance adhered thereto, and

said control device is capable of causing the separate release of said substances from the first and second zones.

5 20. (Original) The stent of claim 19, wherein said substances adhered to said first and second zones are the same substance of the same or a different dose.

 21. (Original) The stent of claim 19, wherein said substances adhered to said first and second zones are different substances.

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22-28. (Canceled)